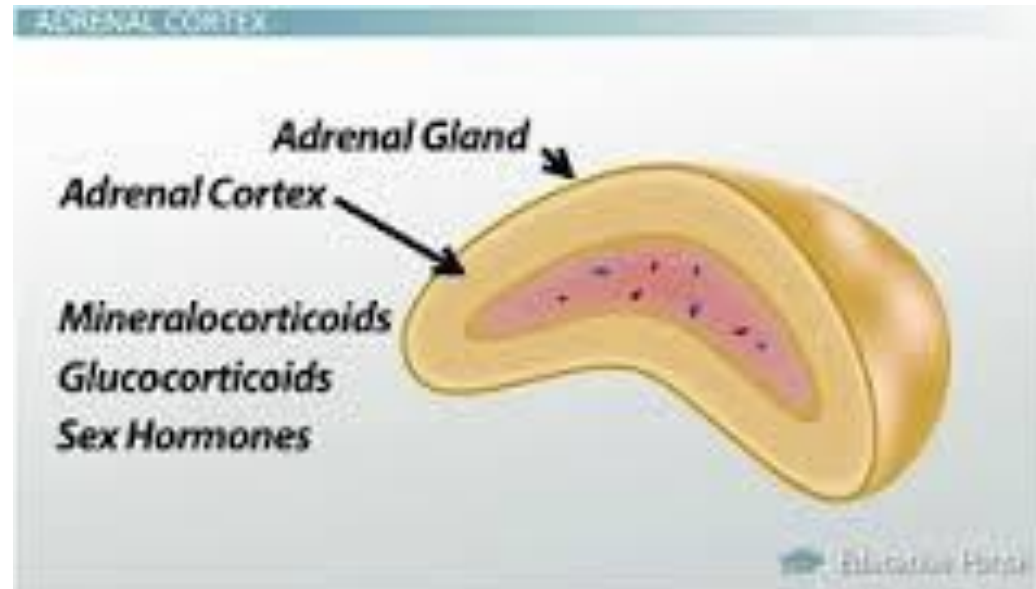


Congenital Adrenal Hyperplasia

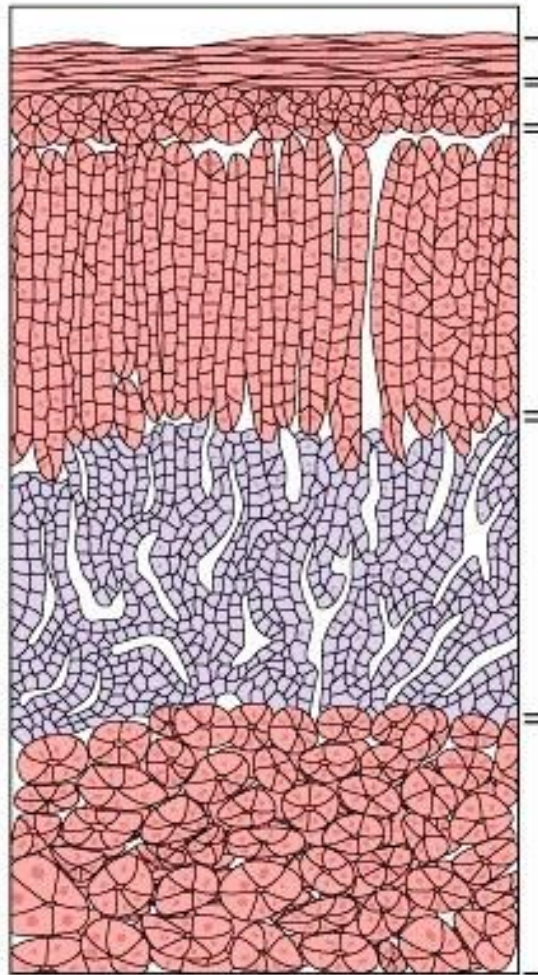
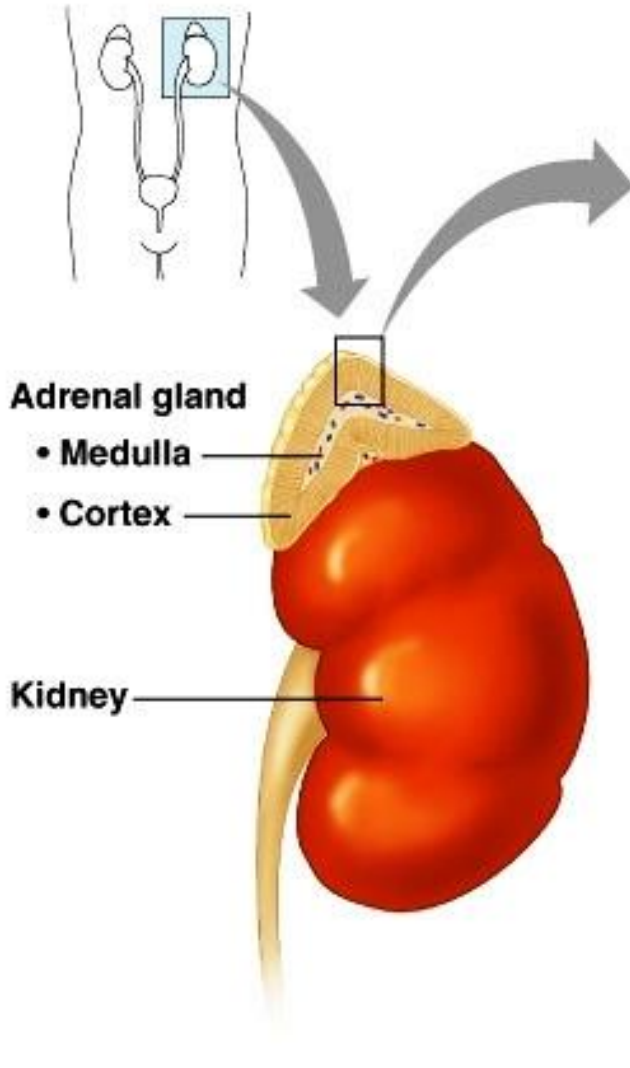


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Objectives

At end of this lecture, students will be able to:

- Recognize the anatomy & physiology of the adrenal gland & the related steroid hormone.
- Recognize the types and presentations of congenital adrenal hyperplasia (CAH).
- Recognize differential diagnosis of different types of CAH and the clinical findings for each.
- Formulate an approach to a child with CAH via history, clinical examination, investigation (electrolyte and acid-base imbalances), and treatment.



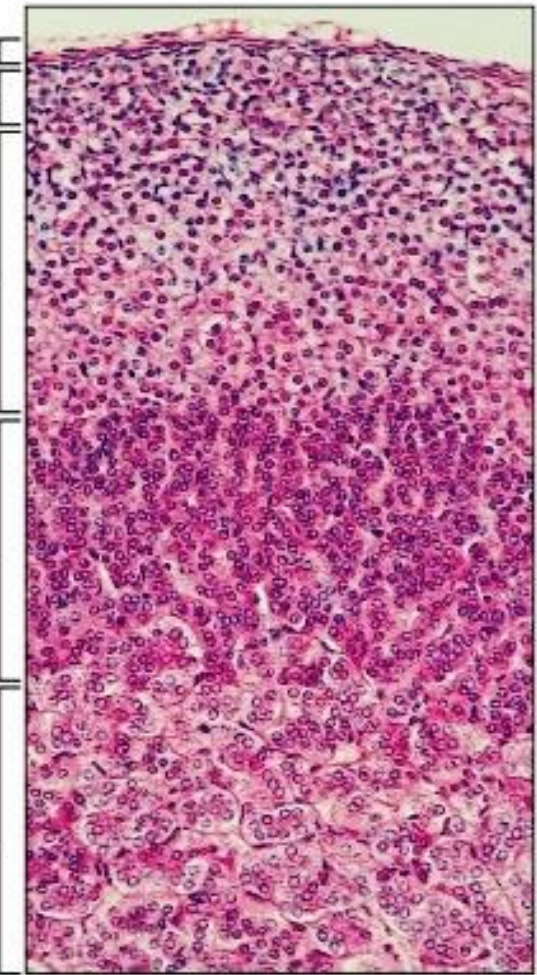
Capsule

Zona glomerulosa

Zona fasciculata

Zona reticularis

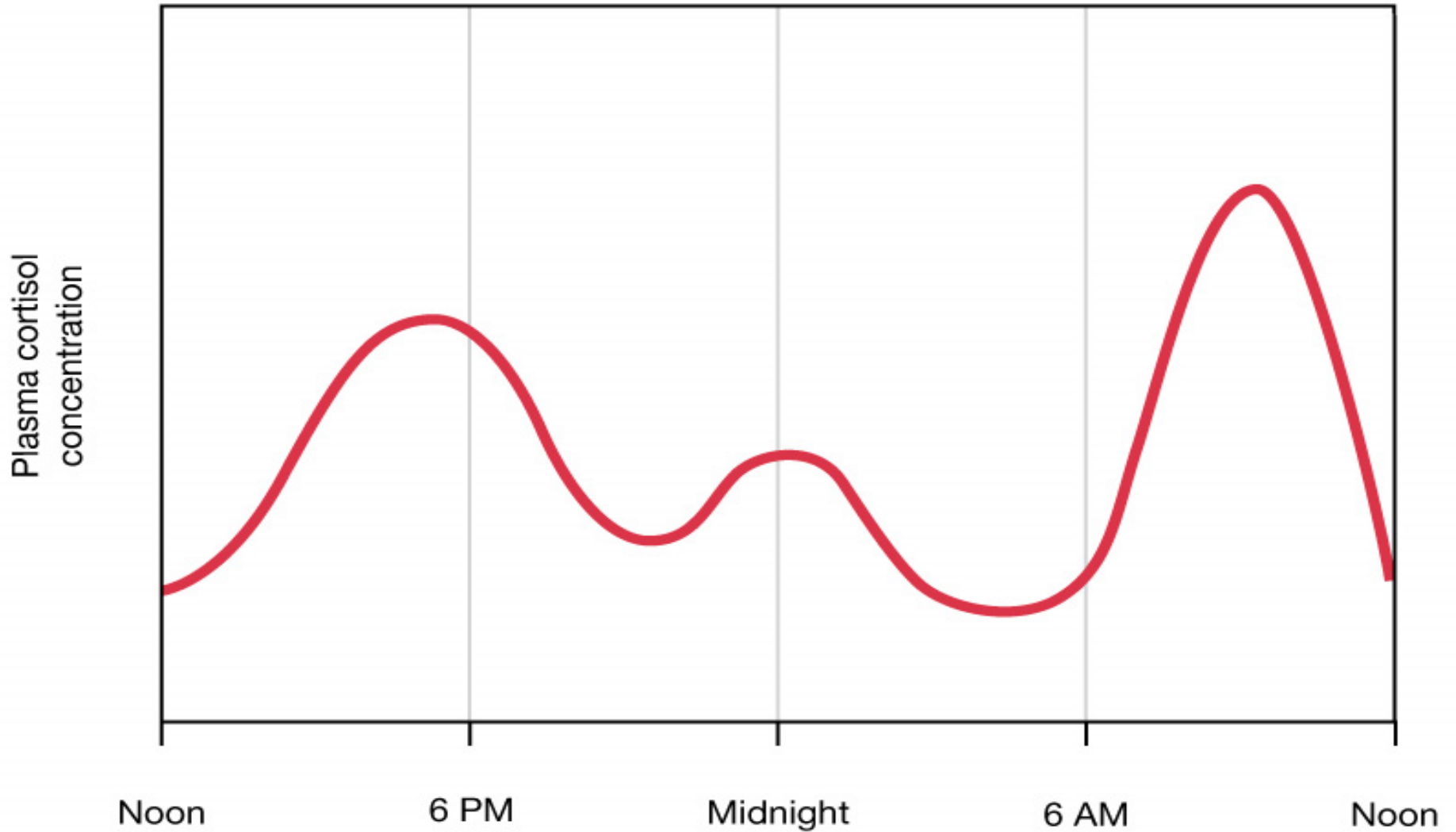
Adrenal medulla

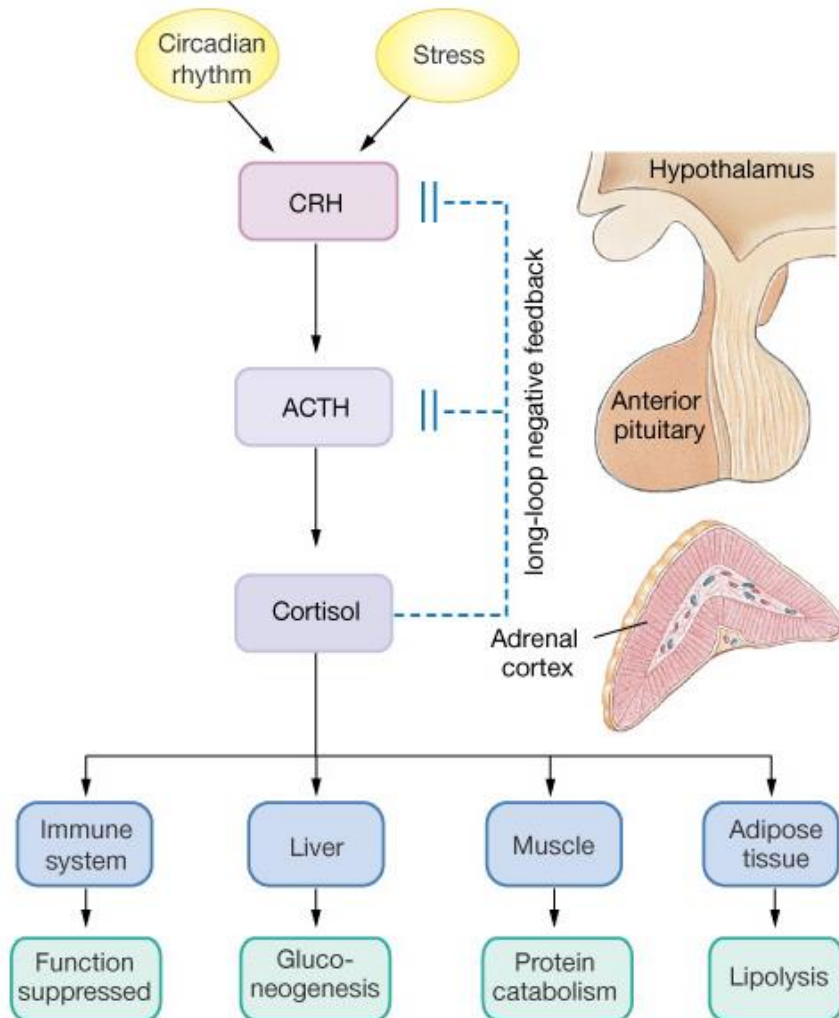


The Adrenal gland

- The adrenal gland lies just above the kidneys
- Divided into two main sub-organs
 - Adrenal cortex
 - Secretes the steroid hormones
 - Glucocorticoid
 - Mineralocorticoid
 - Androgens
 - Adrenal medulla
 - Secretes Catecholamines
 - Adrenaline (epinephrine)
 - Noradrenaline (norepinephrine)

Cortisol Effects: Circadian secretion to match our daily activities





Hypothalamic Pituitary Adrenal Axis

Stimuli

**↓ Blood pressure
↓ Blood flow to kidneys**

**Juxtaglomerular
apparatus in kidneys**

Renin

Angiotensinogen Angiotensin I

ACE

Angiotensin II

Adrenal cortex

Aldosterone

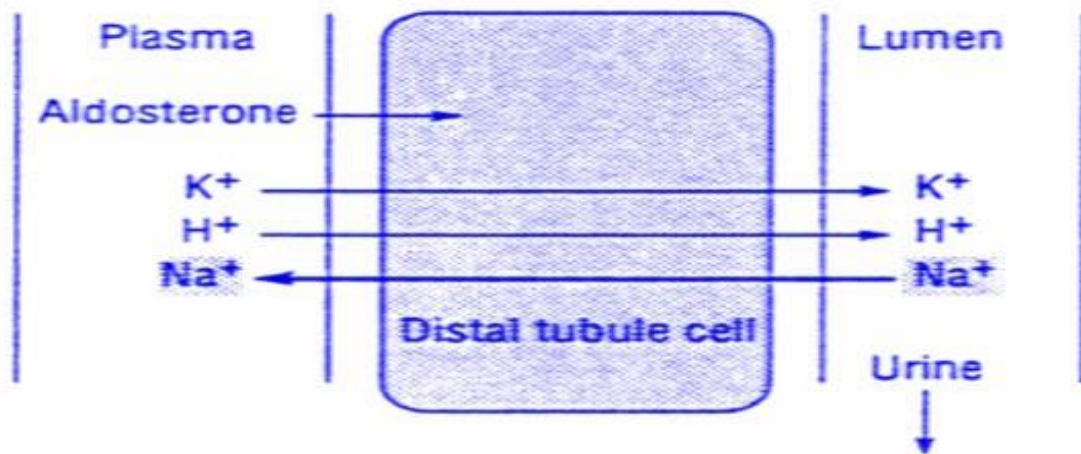
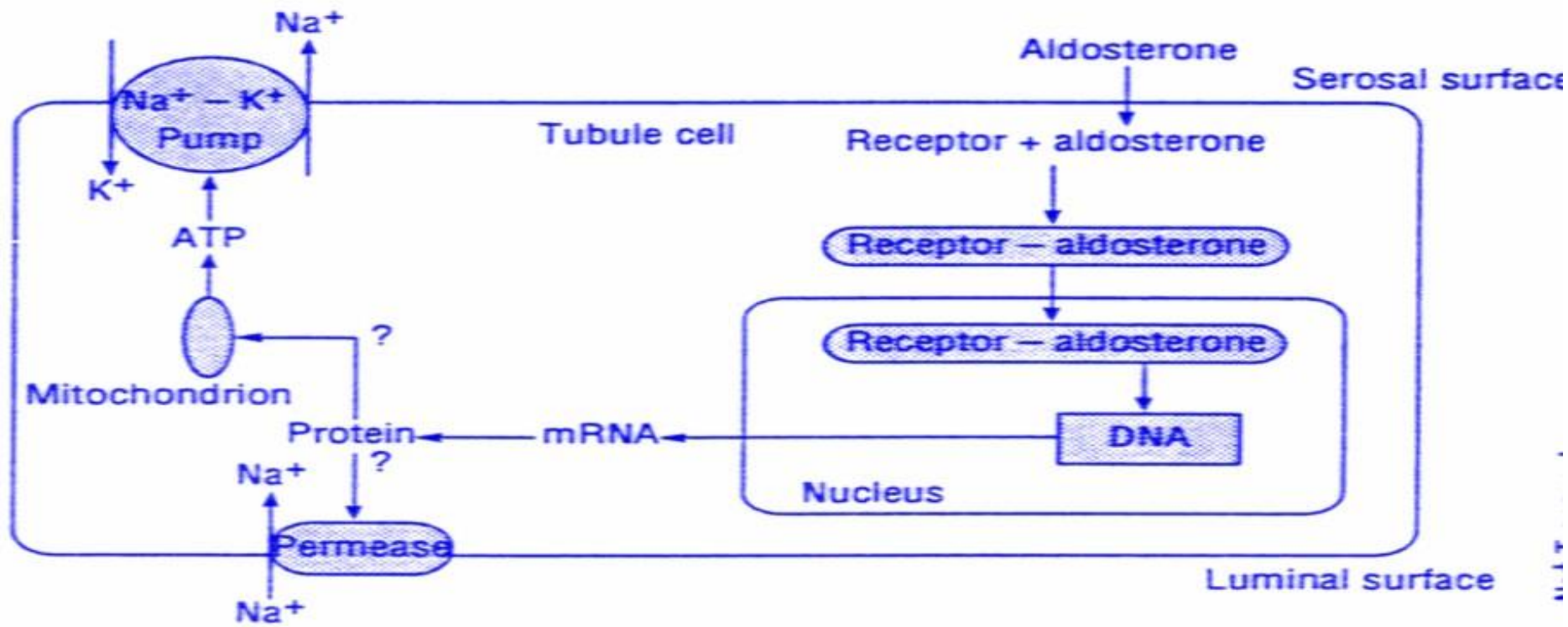
**Salt and water
retention by kidneys**

**Vasoconstriction
of arterioles**

**Negative feedback
responses**

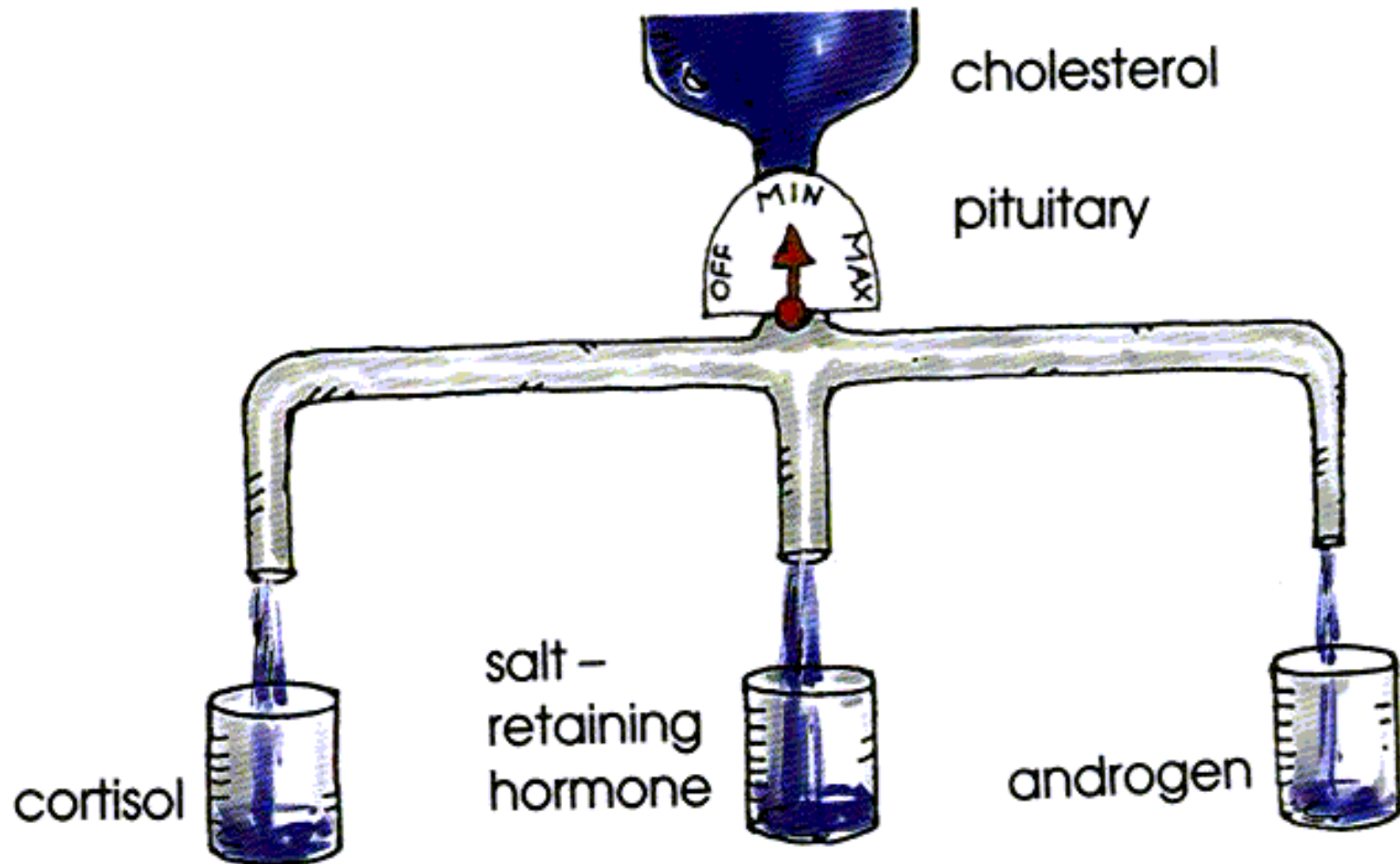
↑ Blood volume

↑ Blood pressure

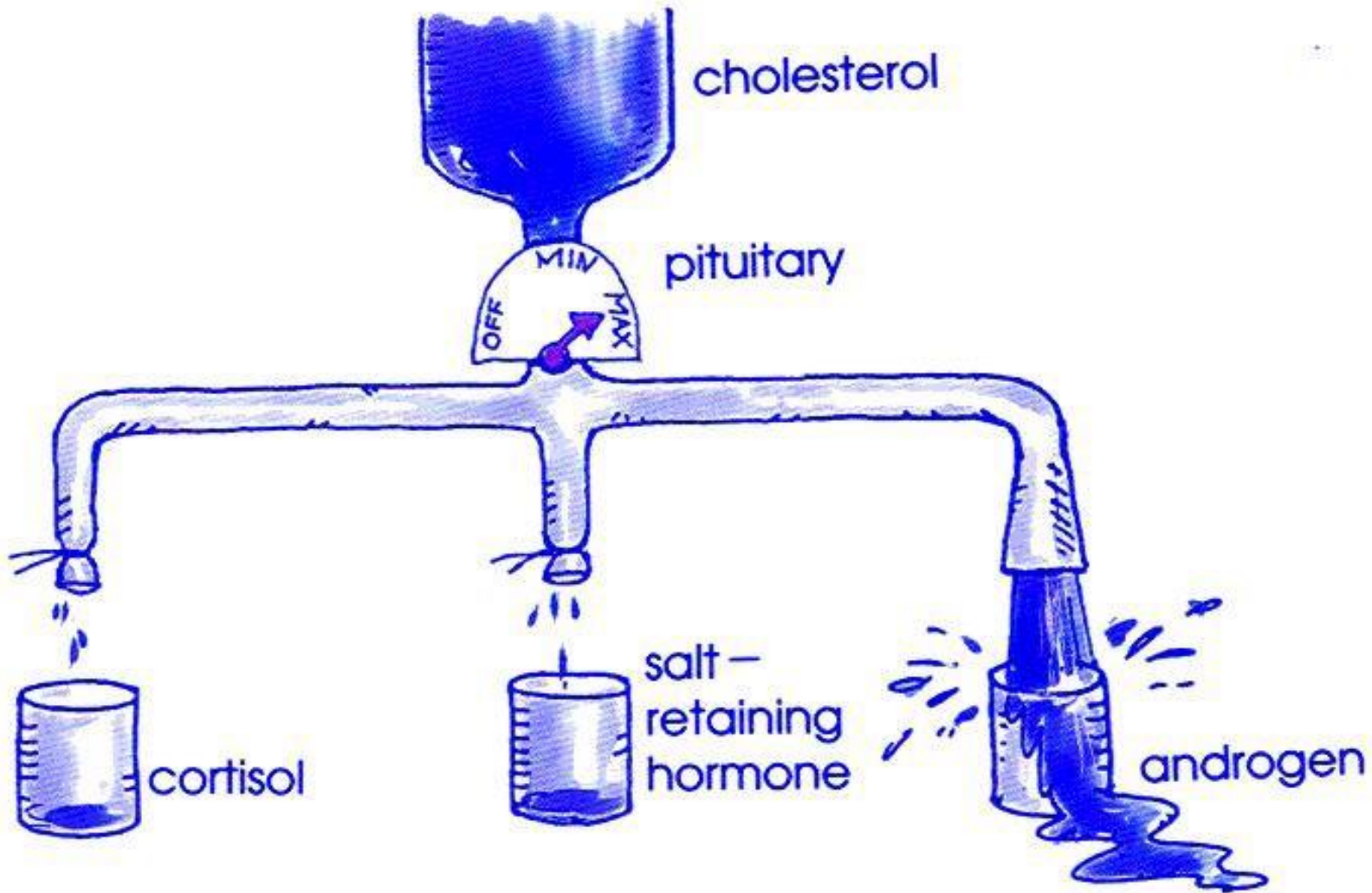


Congenital Adrenal Hyperplasia

Normal Adrenal Cortex Production

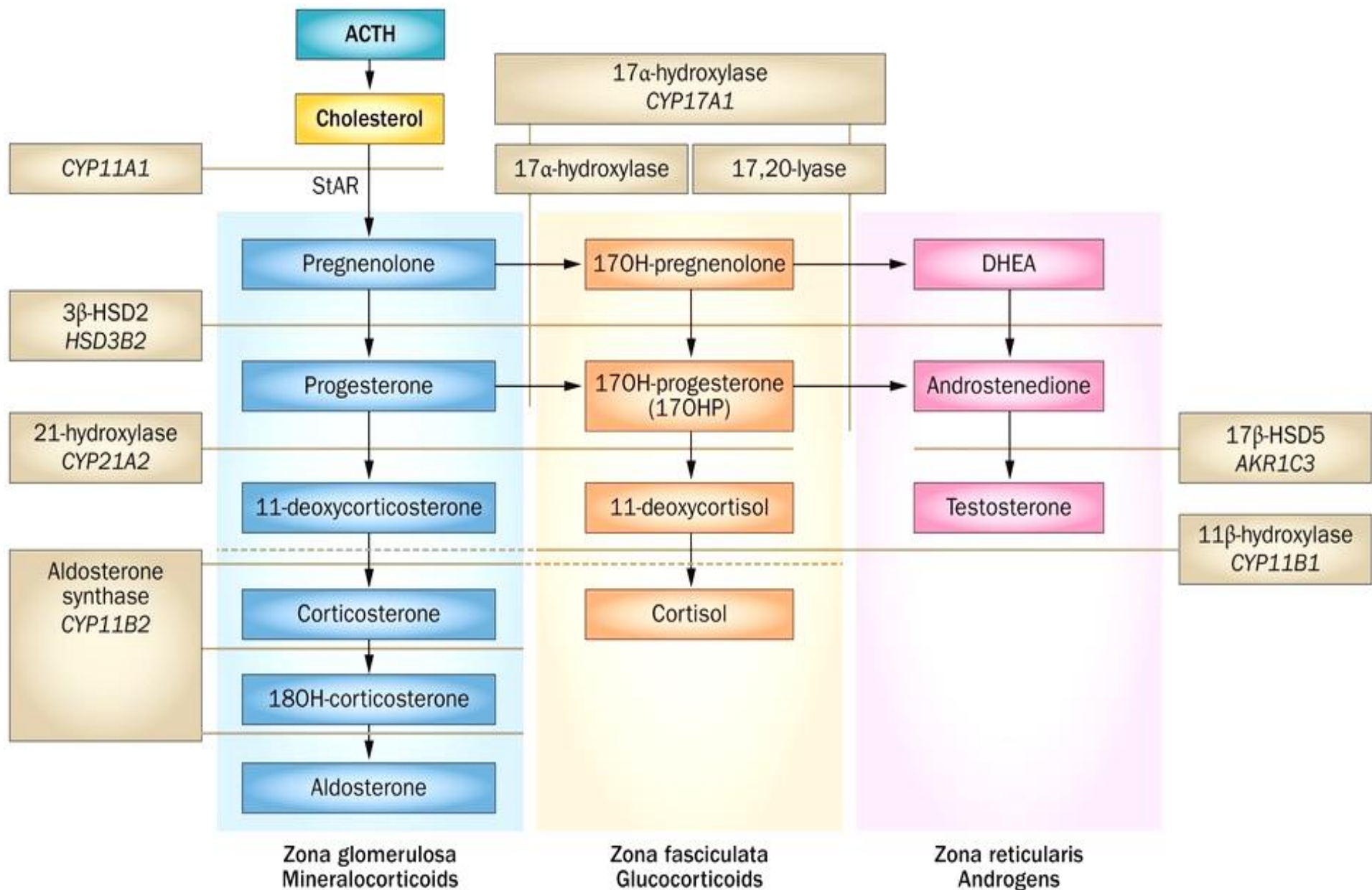


Congenital Adrenal Hyperplasia



Congenital Adrenal Hyperplasia

- It is a familial disorder of adrenal steroid biosynthesis with autosomal recessive mode of inheritance.
- The defect is expressed as adrenal enzyme deficiency.
- 5 major Enzymes deficiency are clinically important but remember only one enzyme is deficient in each patient.
- 90 – 95 % of cases due to 21-Hydroxylase enzyme deficiency.
- Usual age of presentation of classical type is 7-14 days.
 - 21-Hydroxylase enzyme.
 - 11-b-Hydroxylase enzyme.
 - 17-a-Hydroxylase enzyme.
 - 3-b-Hsteroid hydrogenase enzyme.
 - 20,22 Desmolase enzyme.



Congenital Adrenal Hyperplasia

- Autosomal Recessive disease (M=F).
- Incidence 1:1000 -15,000.
- 21-hydroxylase enzyme deficiency is the commonest cause in 90-95 % of cases.
- Gene CYP21 on Chromosome 6
- Neonatal screening on 3rd day of life by doing 17-hydroxyprogesterone (17 OHP).
- Prenatal therapy is effective in preventing genital virilization of affected females.

Congenital Adrenal Hyperplasia

- The clinical phenotype depends upon the nature & severity of the enzyme deficiency.
- Approximately 2/3 of patients with classic CAH due to 21- hydroxylase deficiency have salt wasting due to inadequate Aldosterone synthesis.
- Females are usually recognized at birth because of ambiguous genitalia.

Classical Vs Non- classical CAH

- Depending on the severity of enzyme deficiency, CAH classified into 2 forms:
 - Classical type with moderate – severe enzyme deficiency.
 - Non – classical type with mild enzyme deficiency.
- Classical form presents with early virilization with or without salt-losing crisis, while non-classical type presents with late- onset virilization.
- Non-classical type remains asymptomatic till late childhood when they may show signs of sexual precocity (**precocious puberty**).

Presentations of classical CAH

- Ambiguous genitalia (mainly in females).
- Vomiting and or diarrhea (Gastroenteritis –like).
- Failure to thrive.
- Poor feeding, decreased activity, sleepiness (sepsis –like)
- Dehydration & Shock.
- Salt-wasting presentations with electrolytes imbalance
 - Hyponatremia & hypochloremia
 - Hyperkalemia & metabolic acidosis
- Hypoglycemia.
- Seizure due to low sodium or low glucose.
- Hyperpigmentation due to high ACTH.

Is it a boy or a girl ?





Differential diagnosis of classical CAH

- Neonates with CAH usually presents at age of 7-14 days with:
 - vomiting and electrolyte disturbances so we need to rule out gastroenteritis & pyloric stenosis.
- Poor sucking, poor feeding and sleepiness , need to rule out sepsis.
- Hyponatremia, hyperkalemia with metabolic acidosis, we need to rule out other causes e.g., congenital adrenal hypoplasia, hypoaldosteronism & pseudo hypoaldosteronism .
- Females usually presents with external genital virilization, we need to rule out other causes e.g., maternal androgenic medication intake, maternal ovarian or adrenal tumor during pregnancy and placental aromatase enzyme deficiency.

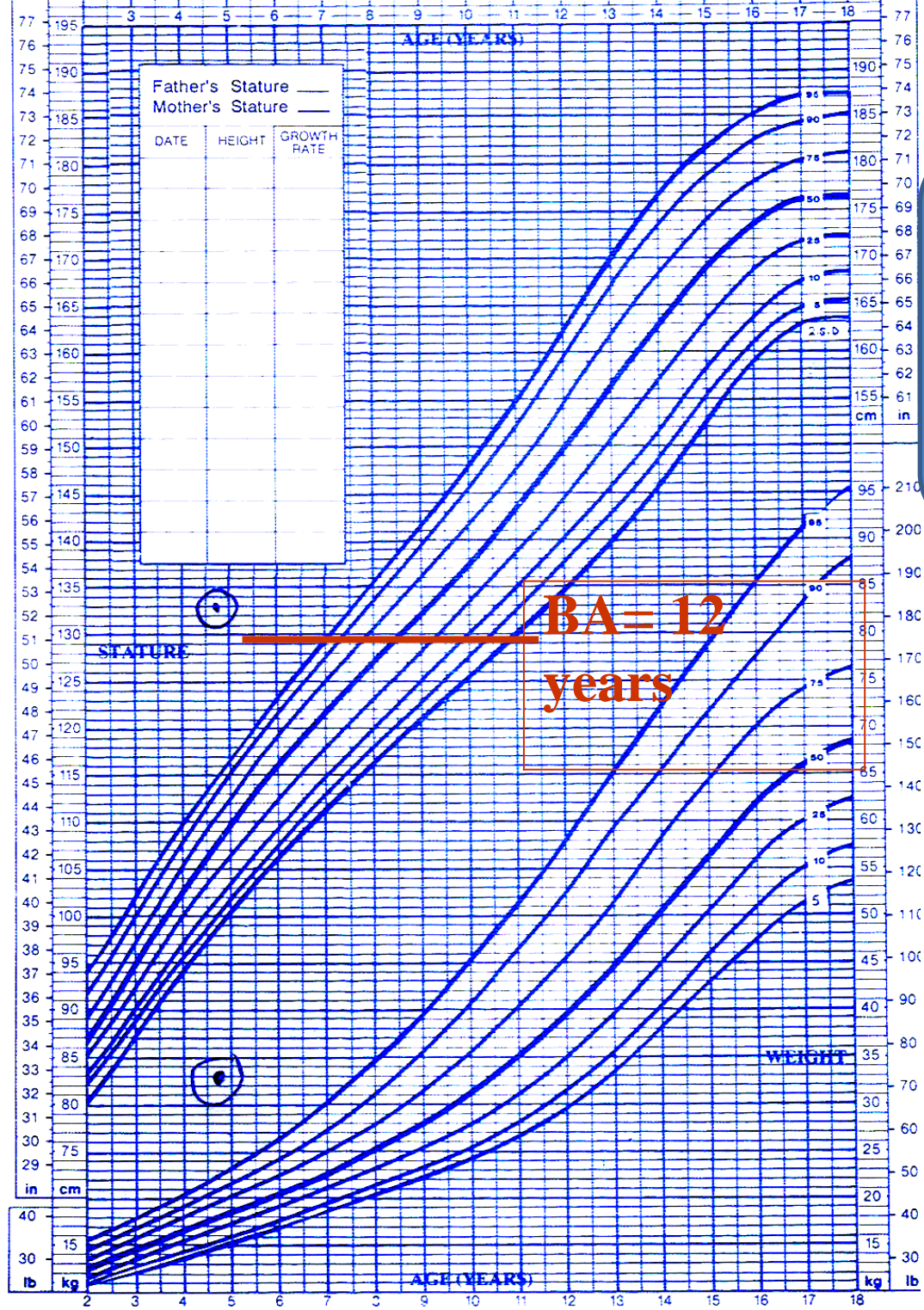
Presentation of Non- Classical CAH

- Precocious puberty.
- Pubarche/ Adrenarche & advanced growth.
- Oligo- Amenorrhea & menstrual irregularity.
- Early beard hair growth.
- Acne.
- Androgenic Alopecia.
- Infertility.
- Need hydrocortisone therapy to suppress adrenal androgens.

Non classical CAH



Six-year-old boy
presented with
peripheral precocious
puberty because of
Non classical CAH




Advance growth with advance bone age in child with non classical CAH

Height = 132.8 cm


Weight = 33 kg



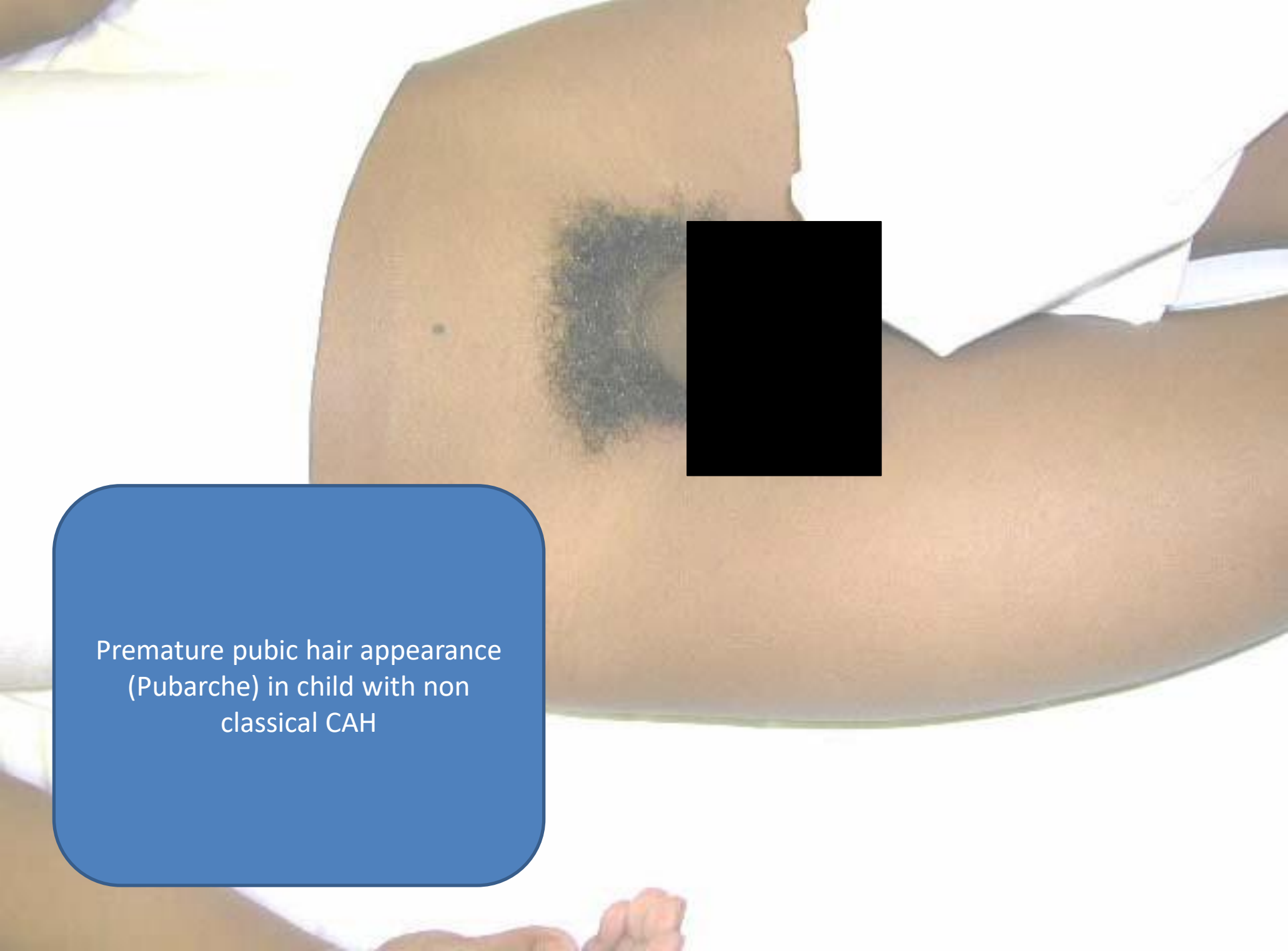
Acne on the back of
non classical CAH

A photograph of a child's legs, showing hyperpigmentation over the knees. The child is lying on a white surface, and the legs are positioned horizontally. The skin is dark brown, and there are distinct, darker patches of hyperpigmentation over the knee joints. A blue rounded rectangle is overlaid on the image, containing white text.

Hyperpigmentation
over the knees in
child with non
classical CAH



Frontal Baldness in
child with non
classical CAH



Premature pubic hair appearance
(Pubarche) in child with non
classical CAH

Differential diagnosis of non classical CAH

- Adrenal tumor.
- Drug-Induced androgen excess.
- Hyperprolactinemia.
- Cushing syndrome.
- Ovarian cancer.
- Ovarian tumor.
- Polycystic ovarian syndrome.

Diagnosis of classical CAH

- A review of a patient's medical history.
- Thorough clinical examination (B.P).
- Serum electrolytes & glucose.
 - Low Na & high K.
 - Hypoglycemia.
 - Elevated serum urea due to associated dehydration.
 - Metabolic acidosis.
- Elevated plasma Renin & ACTH levels.
- Low serum cortisol.
- Low Aldosterone (in salt losing types only).
- High 17 – OHP (screening & confirmation).
- High androgens especially testosterone level.

Management

- Treatment is life-long steroid replacement
 - Hydrocortisone as glucocorticoid agent
 - Fludrocortisone as mineralocorticoid agent.
- Plastic surgery for ambiguous genitalia at early age.
- Genetic counseling.
- Psychological support.

